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(57) Abstract

The gene of the construct pTOM5 is involved in the carotenoid pathway, which produces carotenes, lutein, xanthophylls, and pigments such as lycopene. The invention proposes to modify (inhibit or promote) the synthesis of such compounds in plants using novel DNA constructs comprising a DNA sequence homologous to some or all of the gene encoded by the clone pTOM5, preceded by a plant promoter. In particular, colour of plant parts, especially fruit, may be modified. Yellow tomatoes are disclosed.

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DNA, CONSTRUCTS, CELLS AND PLANTS DERIVED THEREFROM

This application relates to novel DNA constructs, plant cells containing them and plants derived therefrom. In particular it involves the use of recombinant DNA technology to control gene expression in plants.

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In work leading to the present invention we have identified a gene which expresses an enzyme involved in the ripening of tomatoes. This gene has been cloned and characterised. We have now shown that it is involved in carotenoid synthesis. The gene in question is encoded (almost completely) in the clone pTOM5, disclosed by Ray et al (Nucleic Acids Research, 15, 10587, 1989).

The carotenoid biosynthesis pathway is shown 15 in outline in Figure 4 hereof. We believe that the pTOM5 gene is involved in the step or steps of the pathway between geranylgeranyl pyrophosphate and phytoene. Among the products produced by this branch of the pathway are carotenes, lutein, xanthophylls, and pigments such as lycopene. The present invention proposes to modify the synthesis of such compounds in plants.

According to the present invention we provide DNA constructs comprising a DNA sequence homologous to some or all of the gene encoded by the clone pTOM5, preceded by a transcriptional initiation region operative in plants, so that the construct can generate mRNA in plant cells. We further provide a process for modifying the production of carotenoids in plants by transforming such plants

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with constructs according to the present invention. We further provide a process for modifying the production of carotenoids in plants by transforming such plants with DNA adapted to modify carotenoid biosynthesis and growing such transformed plants or their descendants to produce plant parts (for example leaves, petals or fruit) of modified carotenoid content. Suitable DNA comprises, inter alia, constructs according to the present invention, but other similar constructs affecting other parts of the carotenoid pathway may also be used. Such constructs may be adapted to enhance the production of carotenoids (for example lycopene) or inhibit such production by the plant.

As is well known, a cell manufactures protein by transcribing the DNA of the gene for that protein to produce messenger RNA (mRNA), which is then processed (eg by the removal of introns) and finally translated by ribosomes into protein. process may be inhibited by the presence in the cell of "antisense RNA". By this term is meant an RNA sequence which is complementary to a sequence of bases in the mRNA in question: complementary in the sense that each base (or the majority of bases) in the antisense sequence (read in the 3' to 5' sense) is capable of pairing with the corresponding base (G with C, A with U) in the mRNA sequence read in the 5' to 3' sense. It is believed that this inhibition takes place by formation of a complex between the two complementary strands of RNA. preventing the formation of protein. How this works is uncertain: the complex may interfere with further transcription, processing, transport or translation, or degrade the mRNA, or have more than one of these effects. Such antisense RNA may be produced in the cell by transformation with an

appropriate DNA construct arranged to transcribe backwards part of the coding strand (as opposed to the template strand) of the relevant gene (or of a DNA sequence showing substantial homology therewith).

The use of this technology to downregulate the expression of specific plant genes has been described, for example in European Patent publication no 271988 to ICI (corresponding to US serial 119614). Reduction of gene expression has 10 led to a change in the phenotype of the plant: either at the level of gross visible phenotypic difference e.g. lack of anthocyanin production in flower petals of petunia leading to colourless instead of coloured petals (van der Krol et al, 15 Nature, 333, 866-869, 1988); or at a more subtle biochemical level e.g. change in the amount of polygalacturonase and reduction in depolymerisation of pectins during tomato fruit ripening (Smith et al, Nature, 334, 724-726, 1988; Smith et al., Plant 20 Molecular Biology, 14, 369-379, 1990). Thus antisense RNA has been proven to be useful in achieving downregulation of gene expression in plants.

In a further aspect, the present invention provides DNA constructs comprising a transcriptional initiation region operative in plants positioned for transcription of a DNA sequence encoding RNA complementary to a sequence of bases showing substantial homology to an mRNA encoding the enzyme produced by the gene in pTOM5. The invention also includes plant cells containing constructs according to the invention; plants derived therefrom showing modified characteristics, e.g. fruit colour; and fruit and seeds of such plants.

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The constructs of the invention may be inserted into plants to regulate the production of the pTOM5 enzyme. Depending on the nature of the construct, the production of the enzyme may be increased, or reduced, either throughout or at particular stages in the life of the plant. Generally, as would be expected, production of the enzyme is enhanced only by constructs which contain DNA homologous to the substantially complete gene. What is more surprising is that constructs containing an incomplete DNA sequence substantially shorter than that corresponding to the complete gene generally inhibit the expression of the

gene generally inhibit the expression of the enzyme, whether they are arranged to express sense or antisense RNA.

Further evidence for the function of the pTOM5 gene in the carotenoid pathway (se Figure 4) is a significant degree of homology (27% identity; 17% similarity) between the polypeptide predicted from the translation of the pTOM5 sequence and the protein encoded by the crtB gene from Rhodobacter capsulatus, a gram-negative purple non-sulphur bacterium. The crtB gene product catalyses the tail to tail dimerisation of geranylgeranyl pyrophosphate to form prephytoene pyrophosphate. This enzyme (prephytoene synthetase) is the point of divergence of carotenoid biosynthesis from other isoprenoid metabolism. Further, an enzyme has been isolated from Capsicum annuum fruit chromoplasts which is believed to catalyse both the synthesis of prephytoene pyrophosphate and its subsequent conversion to phytoene. This enzyme has a molecular weight of 47,500, in close agreement with the predicted size of the pTOM5 protein (48,000).

As well as colour production, other important functions may be modified by the process of the

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invention. Thus β -carotene (a precursor of Vitamin A) and other carotenoids are important to human health, and have been claimed to have a protective effect against certain diseases. Food plants may be modified by transformation with the constructs of the invention so that they have a higher content of such compounds: or other plants may be so modified, so that they can act as a source from which such compounds can be extracted. Carotenoids are also believed to have a role in protecting plants against high light intensity damage, so plants with a higher content of such compounds may be of value in combating the effects of any global climate change.

. 15 A further possible application of the invention is to control sporopollenin synthesis. This polymer is believed to be a product of carotenoid biosynthesis. Sporopollenin is important during pollen formation and maturation, 20 particularly during the early stages of sporogenesis. Modification of expression of the pTOM5 gene according to the invention in the tapetum is expected to lead to interference in pollen formation, particularly through changing the formation, development or function of the tapetum. 25 Thus inhibition of sporopollenin may be effected in all plants producing sporopollenin through a route involving carotenoids. Accordingly the invention may be used to inhibit the production of pollen in transformed plants. Plants so produced will be 30 male-sterile, and will be useful in the commercial production of uniform hybrids. For this application it is important to use constructs containing promoters which are functional in the tapetum, for example the pollen promoter MFS 14. Not all constitutive promoters work in the tapetum

(for example, the CaMV 35S promoter does not).

The plants to which the present invention can be applied include commercially important fruit-bearing plants, in particular the tomato. In this way, plants can be generated which have modified colour due to promotion or inhibition of the pathways of carotenoid biosynthesis (and in particular of the pigment lycopene).

The invention may be used to promote or inhibit the production of the red colour associated 10 with lycopene. Inhibition of this red colour in tomato fruit (eg by transformation with antisense constructs) gives fruit of an attractive shade of yellow (like certain peppers). Similar yellow tomatoes are known, but the present invention 15 provides a means of transferring the trait into elite lines without a prolonged breeding programme which might alter other traits at the same time. Promotion of lycopene production may produce tomato fruit of a deeper red colour, which may appear more 20 appetising to the consumer, particularly in the form of processed material such as pastes and The invention may also be used to introduce a red colour into parts of plants other than the fruit. Promotion of lycopene may be brought about 25 by inserting one or more functional copies of the gene cDNA, or of the full-length gene, under control of a promoter functional in plants. If lycopene is naturally expressed in the plant, the promoter may 30 be selected to give a higher degree of expression than is given by the natural promoter.

Antisense DNA constructs according to the invention may be very short. They preferably comprise a homologous base sequence at least 10 bases in length. There is no theoretical upper limit to the base sequence — it may be as long as

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the relevant mRNA produced by the cell - but for convenience it will generally be found suitable to use sequences between 100 and 2000 bases in length. The preparation of such constructs is described in more detail below.

The preferred DNA for use in the present invention is DNA derived from the clone pTOM5. required DNA encoding antisense RNA can be obtained in several ways: by cutting with restriction enzymes an appropriate sequence of such DNA; by synthesising a DNA fragment using synthetic oligonucleotides which are annealed and then ligated together in such a way as to give suitable restriction sites at each end; by using synthetic oligonucleotides in a polymerase chain reaction (PCR) to generate the required fragment with suitable restriction sites at each end. The DNA is then cloned into a vector containing upstream promoter and downstream terminator sequences, the cloning being carried out so that the DNA sequence is inverted with respect to its orientation in the strand from which it was cut.

In new vectors expressing antisense RNA, the strand that was formerly the template strand becomes the coding strand, and vice versa. The new vector will thus encode RNA in a base sequence which is complementary to the sequence of pTOM5 mRNA. Thus the two RNA strands are complementary not only in their base sequence but also in their orientations (5' to 3').

As source of the DNA base sequence for transcription, it is convenient to use a cDNA clone such as pTOM5. The base sequence of pTOM5 is set out in Figure 1. This clone has been deposited at the National Collections of Industrial and Marine Bacteria, now at 23 St. Machar Drive, Aberdeen

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AB2 1RY, Scotland, as a plasmid in <u>E.coli</u>, under the reference NCIB 40191, on 1 September 1989. Alternatively, a cDNA clone similar to pTOM5 may be obtained from the mRNA of ripening tomatoes by the method described by Slater et al, Plant Molecular Biology <u>5</u>, 137-147, 1985. In this way may be obtained sequences coding for the whole, or substantially the whole, of the mRNA produced by pTOM5. Suitable lengths of the cDNA so obtained may be cut out for use by means of restriction enzymes.

As previously stated, the preferred source of antisense RNA for use in the present invention is DNA showing homology to the gene encoded by the clone pTOM5 (deposited at the NCIMB as 40191, see above). pTOM5 was derived from a cDNA library isolated from ripe tomato RNA (Slater et al Plant Molecular Biology 5, 137-147, 1985). Three other clones (pTOM45, pTOM91, pTOM104) from the same library cross-hybridise to pTOM5 and probably contain related sequences. pTOM5 has been characterised by hybrid select translation to encode a protein of approximately 48kD (Slater et al, Plant Molecular Biology 5, 137-147, 1985). sequence analysis has demonstrated that the clone is 1600 bases long with an open reading frame encoding a polypeptide of 46.7kD.

We have shown that the mRNA for which pTOM5 codes is expressed in ripening tomato fruit. No expression could be detected in green fruit. pTOM5 is expressed most strongly at the full orange stage of ripening. The level of mRNA then declines in line with the general decline in biosynthetic capacity of the ripening fruit. Expression of pTOM5 mRNA could also be induced by exposing mature green fruit to exogenous ethylene. The expression

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of pTOM5 is reduced in the Ripening inhibitor (\underline{rin}) and Neverripe (\underline{Nr}) tomato fruit ripening mutants which mature very slowly, and never achieve the full red colour associated with ordinary tomato fruit.

The genomic locations in the tomato of sequences homologous to pTOM5 have been identified using RFLP mapping: two loci, on chromosome 2 and chromosome 3 respectively, carry sequences homologous to pTOM5. It has also been shown by Southern blotting that the gene may be present as a small multigene family. Genomic clones representing two individual genes have been isolated and characterised by DNA sequence analysis. gTOM5 represents represents a gene with exon sequence identical to pTOM5. Clone F contains sequence similar, but not identical to pTOM5. Sequence and expression data suggest that Clone F encodes an untranscribed pseudogene.

Although there is a considerable body of information on the structure and expression of the pTOM5 gene family, the biochemical function of this clone has not hitherto been known.

An alternative source of DNA for the base sequence for transcription is a suitable gene encoding the pTOM5 enzyme. This gene may differ from the cDNA of pTOM5 in that introns may be present. The introns are not transcribed into mRNA (or, if so transcribed, are subsequently cut out). When using such a gene as the source of a partial base sequence for transcription it is possible to use either intron or exon regions.

A further way of obtaining a suitable DNA base sequence for transcription is to synthesise it <u>ab</u> <u>initio</u> from the appropriate bases, for example using Figure 1 as a guide.

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Recombinant DNA and vectors according to the present invention may be made as follows. A suitable vector containing the desired base sequence for transcription (for example pTOM5) is treated with restriction enzymes to cut the sequence out. The DNA strand so obtained is cloned (if desired, in reverse orientation) into a second vector containing the desired promoter sequence, for example cauliflower mosaic virus 35S promoter or the tomato polygalacturonase gene promoter sequence — Bird et al., Plant Molecular Biology, 11, 651-662, 1988) and the desired terminator sequence (for example the 3' of the Agrobacterium tumefaciens nopaline synthase gene, the nos 3' end).

According to the invention we propose to use both constitutive promoters (such as cauliflower mosaic virus 35s) and inducible or developmentally regulated promoters (such as the ripe-fruit 20 specific polygalacturonase promoter) as circumstances require. Use of a constitutive promoter will tend to affect functions in all parts of the plant: while by using a tissue-specific promoter, functions may be controlled more selectively. Thus in applying the invention, e.g. 25 to tomatoes, it may be found convenient to use the promoter of the PG gene (Bird et al, 1988, cited above). Use of this promoter, at least in tomatoes, has the advantage that the production of recombinant RNA is under the control of a ripening-. 30 specific promoter. Thus the recombinant RNA is only produced in the organ in which its action is required. By this means particular organs can have their colour modified. For plants to be used as ornament (as many are), by using a constitutive promoter one may induce carotenoid production

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throughout the plant to provide plants with an overall modified or reddish/yellowish colour: normally it will be preferred to limit the production to particular sites (such as fruit, flower petals or tubers) by using a tissue-specific or developmentally regulated promoter, for example the PG gene promoter referred to above. Other ripening-specific promoters that could be used include the E8 promoter (Diekman & Fischer, EMBO Journal 7, 3315-3320, 1988) and the 2A11 promoter from tomatoes described in US Patent 4,943,674 to Calgene.

Vectors according to the invention may be used to transform plants as desired, to make plants according to the invention. Dicotyledonous plants, such as tomato, may be transformed by Agrobacterium Ti plasmid technology, for example as described by Bevan (1984) Nucleic Acid Research, 12, 8711-8721. Such transformed plants may be reproduced sexually, or by cell or tissue culture.

The degree of production of RNA in the plant cells can be controlled by suitable choice of promoter sequences, or by selecting the number of copies, or the site of integration, of the DNA sequences according to the invention that are introduced into the plant genome. In this way it may be possible to modify colour formation to a greater or lesser extent.

The constructs of our invention may be used to transform cells of both monocotyledonous and dicotyledonous plants in various ways known to the art. In many cases such plant cells (particularly when they are cells of dicotyledonous plants) may be cultured to regenerate whole plants which subsequently reproduce to give successive generations of genetically modified plants.

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Examples of genetically modified plants according to the present invention include, as well as tomatoes, fruits of such as mangoes, peaches, apples, pears, strawberries, bananas and melons. The fruit of such plants may be made more attractive (or at least interesting) by inducing or intensifying a red colour therein. Other plants that may be modified by the process of the invention include tubers such as radishes, turnips and potatoes, as well as cereals such as maize (corn), wheat, barley and rice. Flowers of modified colour, and ornamental grasses either red or reddish overall, or having red seedheads, may be produced.

15 Plants produced by the process of the invention may also contain other recombinant constructs, for example constructs having other effects on fruit ripening. In particular tomatoes of enhanced colour according to the invention may 20 also contain constructs inhibiting the production of enzymes such as polygalacturonase and pectinesterase, or interfering with ethylene production (eg from pTOM13, see PCT Application 90/01072 filed 12 July 1990). Such tomatoes can have higher solids contents than conventional 25 tomatoes and produce more tomato paste per unit of fruit weight. The extra lycopene production in such tomatoes is desirable to prevent any lightening of colour that might otherwise be observed in such pastes. Tomatoes containing both 30 types of recombinant construct may be made either by successive transformations, or by crossing two varieties that each contain one of the constructs, and selecting among the progeny for those that contain both.

The invention will now be described further

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with reference to the accompanying drawings, in which:

Figure 1 shows the base sequence of the clone pTOM5;

Figure 2 shows the regions of the pTOM5 sequence which may be synthesised by polymerase chain reaction and used in the construction of vectors according to the invention;

Figure 3 shows the base sequence of the oligonucleotides used as primers for the polymerase chain reactions to synthesise the fragments illustrated in Figure 2;

Figure 4 is a diagram of the carotenoid synthesis pathway;

Figure 5 is a diagram of DNA vector pCB17; Figure 6 is a diagram of DNA vector pCB19;

Figure 7 diagrams the scheme for the manufacture of construct pJREX5;

Figure 8 diagrams the scheme for the 20 manufacture of construct pCBEX5.

The following Examples illustrate aspects of the invention.

25 EXAMPLE 1

Construction of pTOM5 antisense RNA vectors with the CaMV 35S promoter ${\bf r}$

Three vectors may be constructed using sequences corresponding to different lengths of the pTOM5 cDNA as shown in figure 2.

- 1. bases 1 to 187 pJR15A
- 2. bases 1 to 794 pJR15B
- 3. bases 1 to 1598 (the complete cDNA) pJR15C

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pJR15B was synthesised in vitro using polymerase chain reactions with synthetic oligonucleotides T5AS-1 and T5AS-3 shown in Figure 2 as primers and pTOM5 cDNA as template. synthetic oligo nucleotide primers were designed such that a BamHI restriction site was incorporated at the 5' end a KpnI site was incorporated at the 3' end of the fragment. After cleavage with BamHI and KpnI, the fragment was cloned into the vector pJR1 which had previously been cut with KpnI and BamHI. pJR1 (Smith et al Nature 334, 724-726, 1988) is a Bin19 (Bevan, Nucleic Acids Research, 12, 8711-8721, 1984) based vector, which permits the expression of the antisense RNA under the control of the CaMV 35S promoter. This vector includes a nopaline synthase (nos) 3' end termination sequence.

After synthesis of the vector pJR15B, the structures and orientation of the pTOM5 sequence was confirmed by DNA sequence analysis.

Vectors pJR15A and pJR15C are made similarly, following the construction schemes shown in Figures 2 and 3.

25 EXAMPLE 2

The fragments of the pTOM5 cDNA described in Example 1 are also cloned into the vector pJR2 to give the following clones:

bases 1 to 187 - pJR25A
 bases 1 to 794 - pJR25B
 bases 1 to 1598 - pJR25C

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pJR2 is a Bin19 based vector, which permits the expression of the antisense RNA under the control of the tomato polygalacturonase promoter. This vector includes a nopaline synthase (nos) 3' end termination sequence. This vector does not contain a KpnI or a BamHI site between the promoter and terminator sequences. Consequently, the PCR synthesised fragments are digested with KpnI and BamHI, the cut ends are made flush with T4 polymerase and then cloned into the HincII site of pJR2

After synthesis, vectors with the correct orientation of pTOM5 sequence are identified by DNA sequence analysis.

15 EXAMPLE 3

Construction of pTOM5 sense RNA vectors with the CaMV 35S promoter $\ \ \,$

The fragments of pTOM5 cDNA described in Example 2 are also cloned into the vector pJR1 in the sense orientation to give the following clones:

bases 1 to 187 - pJR15AS
 bases 1 to 794 - pJR15BS
 bases 1 to 1598 - pJR15CS

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The PCR generated fragments are digested with KpnI and BamHI, the cut ends made flush with T4 polymerase and then cloned into the HincII site of pJR1. After synthesis, the vectors with the sense orientation of pTOM5 sequence are identified by DNA sequence analysis.

EXAMPLE 4

Generation and analysis of plants transformed with the vector pJR15B.

The pJR15B vector was transferred to Agrobacterium tumefaciens LBA4404 (a micro-organism

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widely available to plant biotechnologists) and used to transform tomato plants. Transformation of tomato stem segments followed standard protocols (e.g. Bird et al Plant Molecular Biology 11, 651-662, 1988). Transformed plants were identified by their ability to grow on media containing the antibiotic kanamycin. Forty-one individual plants were regenerated and grown to maturity. Thirty-seven of these plants produced fruit which changed colour to yellow rather than to red: they did not turn red even when over-ripe. Fruit from the other four plants turned orange-red.

The flowers of plants with yellow fruit had pale corollae. The accumulation of yellow pigment in flowers varied between individual transformants with some flowers being almost white.

Preliminary analysis indicated that carotenoid accumulation in the yellow fruit was approximately 6% of that in untransformed controls. Almost no lycopene was detected (<2% of that in normal fruit): the majority of residual carotenoid was lutein and β -carotene, neither of which accumulated to significantly greater levels than in the control fruit. Most of the yellow fruit pigment was in the skin, and could not be extracted by methanol. Thus it is unlikely to be carotenoid.

Three of the transformants giving yellow fruit and two of the transformants giving red fruit were submitted to polymerase chain reaction analysis. This indicated that the pJR15B antisense construct was present and intact in all five. DNA blot analysis indicated that the insert copy number was between 1 and 4.

One transformed plant, coded E64C8, with yellow fruit, was selfed to produce progeny. These showed segregation of yellow and white flower

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colour according to Mendelian principles, indicating that the plant contains one functional copy of the antisense construct, and that the phenotype is stably inherited.

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EXAMPLE 5

Construction of pTOM5 expression vectors with the CaMV 35S promoter.

Construction of pJREX5

The expression vector pJREX5 is synthesised in vitro according to the scheme shown in Figure 7.

The 1468 bp SspI fragment is isolated from pTOM5, the cut ends are made flush with T4 polymerase and the resulting product is then cloned into the SmaI site of the plasmid pJR1.

After synthesis, vectors with sense and antisense orientation of the pTOM5 fragment are identified by DNA sequence analysis.

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EXAMPLE 6

Construction of pTOM5 expression vectors with the tomato polygalacturonase promoter.

A. Construction of pCB17.

A 1.6 kb region from the 3' end of the tomato PG gene was substituted for the nopaline synthase polyadenylation sequence in pCB1 (Bird et al Plant Molecular Biology 11, 651-662, 1988).

The 5.8 kb Sall/BamHI fragment adjacent to the right arm of lambda EMBL3 in gTOM23 was cloned into the Sall/BamHI sites of pUC8 to give plasmid pGTOM23.5.8. The 1.6 kb BgIII fragment from pGTOM23.5.8 was isolated and cloned into the BamHI site of pUC19. Plasmids with the correct orientation of the 1.6 kb BgIII insert contained a 550 bp Xbal/BstEII fragment. One such clone was designated A3/1.

A 2.2 kb HindIII/PvuI was isolated as a fragment from pCB1. This contained a 1.45kb PG promoter fragment and the chloramphenicol acetyl transferase (CAT) gene. This was cloned into Bin19 (Bevan, Nucleic Acids Research, 1984, 12, 5 8711-8721) which had been cut with SalI followed by filling of the cohesive ends with T4 DNA polymerase and subsequently digested with HindIII. Plasmids with the 2.2 kb HindIII/PvuI fragment contained a 2.2 kb HindIII/XbaI fragment. One of these clones 10 was digested with XbaI and KpnI and ligated with the 1.6 kb XbaI/KpnI fragment from A3/1. After transformation, one clone with the correct insert was designated pCB17 (Figure 5).

15 The correct construction of pCB17 was checked by nucleotide sequence analysis of the plasmid DNA at the boundary between the CAT gene and the PG 3' fragment. An unexpected region of the Bin19 polylinker was found to have remained at this junction. This was judged to be unlikely to interfere with the correct functioning of the plasmid. The sequence of pCB17 in this region is:

25 CAT BIN19 POLYLINKER PG 3'
...CCGTCCCCGTGCATGCCTGCAGGTCGACTCTAGAGGATCTTCAATATA
TAG... Xbai

B. Construction of pCB19

The PG promoter region in plasmid pCB17 was extended by the addition of a 3.5 kb fragment from genomic clone gTOM23 (NCIMB Accession Number 12373).

The 3.5 kb HindIII fragment from pGTOM23.7.8 was cloned into the HindIII site in pCB17. Plasmids with the correct orientation of the 3.5 kb

Hind AYE insert contained a 4.9 kb SaII/BamI fragment. One such clone was designated pCB19 (Figure 6).

5 C. Construction of pCB19

The expression vector pCBEX5 is synthesised in vitro according to the scheme shown in Figure 8. The 1468 bp SspI fragment is isolated from pTOM5, and the cut ends are made flush with T4 polymerase. The resulting product is then cloned into pCB19 from which the BamHI - XbaI fragment encoding chloramphenicol acetyl transferase has been deleted

After synthesis, vectors with sense and antisense orientation of the pTOM5 fragment are identified by DNA sequence analysis.

and the cut ends made flush with T4 polymerase.

EXAMPLE 7

Generation of transformed plants with the 20 sense vectors pJREX5 and pCBEX5. The required sense expression vector (produced in Example 5 or 6) is transferred to Agrobacterium tumefaciens LBA4404 (a micro-organism widely available to plant biotechnologists) and used to 25 transform tomato plants. Transformation of tomato stem segments follow standard protocols (e.g. Bird et al Plant Molecular Biology 11, 651-662, 1988). Transformed plants were identified by their ability to grow on media containing the antibiotic 30 kanamycin and by the detection of DNA by DNA blot analysis of their genomic DNA. Ripening fruit are analysed for levels of lycopene and other carotenoids. Plants with higher than normal lycopene levels are selected for further use and study.

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CLAIMS:

- 1. DNA constructs comprising a DNA sequence showing homology to at least some of the gene for the gene product encoded by the clone pTOM5, preceded by a transcriptional initiation region operative in plants, so that the construct can generate mRNA in plant cells.
- DNA constructs as claimed in claim 1 comprising a transcriptional initiation region
 operative in plants positioned for transcription of a DNA sequence encoding RNA complementary to a base sequence showing substantial homology to an mRNA encoding the enzyme produced by the gene of pTOM 5.
- 3. DNA constructs as claimed in claim 1 comprising a transcriptional initiation region operative in plants positioned for transcription of a DNA sequence to give mRNA which can be translated to give the enzyme produced by the gene of pTOM 5.
 - 4. DNA constructs as claimed in claim 1 comprising a transcriptional initiation region operative in plants positioned for transcription of a DNA sequence encoding RNA showing substantial homology to part only of the mRNA encoding the enzyme produced by the gene of pTOM 5.
 - 5. DNA constructs as claimed in any of claims 1 to 4 in which the transcriptional initiation region is a constitutive promoter such as CaMV 35s.
 - 6. DNA constructs as claimed in any of claims 1 to 5 in which the transcriptional initiation region is an inducible or developmentally-regulated or

tissue-specific promoter.

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- 7. A process for modifying the production of carotenoids in plants by transforming such plants with DNA adapted to modify carotenoid biosynthesis and growing such transformed plants or their descendants to produce plant parts of modified carotenoid content.
- 8. Process as claimed in claim 7 in which the DNA is a construct claimed in any of claims 1 to 6.

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- 9. A process for producing plants having parts of modified colour which comprises transforming plants with DNA adapted to modify production of lycopene therein and growing such transformed plants or
- 20 their descendants to produce plant parts of changed colour.
- 10. Process as claimed in claim 9 in which the plant naturally produces lycopene and the construct25 is adapted to inhibit the production of lycopene.
 - 11. Process as claimed in claim 10 in which the DNA construct is one claimed in either of claims 2 or 4.

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- 12. Process as claimed in claim 9 for producing or intensifying a red colour in plant parts in which the construct is adapted to promote the construction of lycopene.
- 13. Process as claimed in any of claims 10 to 12 in which the construct is one claimed in claim 3.

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- 14. Process as claimed in any of claims 7-13 inwhich the plant is a tomato.
 - 15. Transformed plants produced in the processes of any of claims 7-13.
- 16. Plants as claimed in claim 15 which are fruit, eg tomatoes, mangoes, peaches, apples, pears, strawberries, bananas, melons or citrus fruit.
- 15 17. Plants as claimed in claim 15 which are tubers such as radishes, turnips and potatoes.
 - 18. Plants as claimed in claim 15 which are cereals such as maize (corn), wheat, barley and rice.
 - 19. Plants as claimed in claim 15 that are ornamental in function, e.g. flowers.
- 25 20. Plant parts of modified colour harvested from plants claimed in any of claims 15-19.
 - 21. Fruit and seed of plants claimed in any of claims 15-19.
 - 22. A tomato as claimed in claim 14 which additionally comprises a recombinant construct inhibiting the production of polygalacturonase or pectinmethylesterase.

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FIG. 1

SEQ ID NO: 1

SEQUENCE TYPE: Nucleotide SEQUENCE LENGTH: 1646 base pairs

STRANDEDNESS: single TOPOLOGY: linear MOLECULE TYPE: CDNA

ORIGINAL SOURCE ORGANISM: Tomato var. Ailsa Craig IMMEDIATE EXPERIMENTAL SOURCE: Ripe fruit cDNA library

FEATURES:

from 201 to 1436 bp open reading frame

PROPERTIES: cDNA of tomato fruit ripening related gene - pTOM5

TTTGCCTGTC		TTATAATCTT	TTTCTACAGA	AGAGAAAGTG	GGTAATTTTG	60
TTTGAGAGTG		CTAGTGGGAA				
TAAGTAAAGT		ACAAAAAGAA	AGACAAAAT	CTTGGAATTG		
CCAAGGTTTT	CTTGCTCAGA	ATGTCTGTTG	CCTTGTTATG	GGTTGTTTCT		
TCTCAAATGG	GACAAGTTTC	ATGGAATCAG	TCCGGGAGGG		00110100	
CGAGGCATAG	GAATTTGGTG	TCCAATGAGA	GAATCAATAG	AGGTGGTGGA		
ATAATGGACG		GTACGGTCTG	CTATTTTGGC	TACTCCATCT	GGAGAACGGA	
CGATGACATC		GTCTATGATG	TGGTTTTGAG	GCAGGCAGCC	TTGGTGAAGA	
GGCAACTGAG	ATCTACCAAT	GAGTTAGAAG	TGAAGCCGGA	TATACCTATT		540
TGGGCTTGTT	GAGTGAAGCA	TATGATAGGT	GTGGTGAAGT	ATGTGCAGAG	TATGCAAAGA	600
CGTTTAACTT	AGGAACTATG	CTAATGACTC	CCGAGAGAAG	AAGGGCTATC	TGGGCAATAT	660
ATGTATGGTG	CAGAAGAACA	GATGAACTTG	TTGATGGCCC	AAACGCATCA		720
CGGCAGCCTT		GAAAATAGGC	TAGAAGATGT	TTTCAATGGG	CGGCCATTTG	780
ACATGCTCGA	TGGTGCTTTG	TCCGATACAG	TTTCTAACTT	TCCAGTTGAT	ATTCAGCCAT	840
TCAGAGATAT	GATTGAAGGA	ATGCGTATGG	ACTTGAGAAA	ATCGAGATAC	AAAAACTTCG	900
ACGAACTATA	CCTTTATTGT	TATTATGTTG	CTGGTACGGT	TGGGTTGATG	AGTGTTCCAA	960
TTATGGGTAT	CGCCCCTGAA	TCAAAGGCAA	CAACAGAGAG	CGTATATAAT	GCTGCTTTGG	1020
CTCTGGGGAT	CGCAAATCAA	TTAACTAACA	TACTCAGAGA	TGTTGGAGAA	GATGCCAGAA	1080
GAGGAAGAGT	CTACTTGCCT	CAAGATGAAT	TAGCACAGGC	AGGTCTATCC	GATGAAGATA	1140
TATTTGCTGG	AAGGGTGACC	GATAAATGGA	GAATCTTTAT	GAAGAAACAA		1200
CAAGAAAGTT	CTTTGATGAG	GCAGAGAAAG	GCGTGACAGA	ATTGAGCTCA	GCTAGTAGAT	1260
TCCCTGTATG	GGCATCTTTG	GTCTTGTACC	GCAAAATACT	AGATGAGATT	GAAGCCAATG	1320
ACTACAACAA	CTTCACAAAG	AGAGCATATG	TGAGCAAATC	AAAGCAAGTT	GATTGCATTA	1380
CCTATTGCAT	ATGCAAAATC	TCTTGTGCCT	CCTACAAAAC	TGCCTCTCTT	CAAAGATAAA	1440
GCATGAAATG	AAGATATATA	TATATATATA	TATAGCAATG	TACATTAGAA	GAAAAAAAGG	1500
AAGAAGAAAT	GTTGTTGTAT	TGATATAAAT	GTATATCATA	AATATTAGGT	TGTAGTAACA	1560
TTCAATATAA	TTATCTCTTG	TAGTTGTTGT	ATCTTCACTT	TATCTCAACT	CCTTTGAGAG	1620
AACTTTCCGT	AAAAAAAAA	AAAAA		CICANCI	CCITIGAGAG	1646
						TD40

Construct pJR15A pJR15B pJR15C Strategy for synthesis of BamHI-KpnI fragments pTOM5 1600 bp PCR oligos by PCR from pTOM5 F16.2 T5AS-3 Kpn **▲** T5AS-2 T5AS-1 BamHI BamHI Fragment 4 m ပ

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FIG. 3

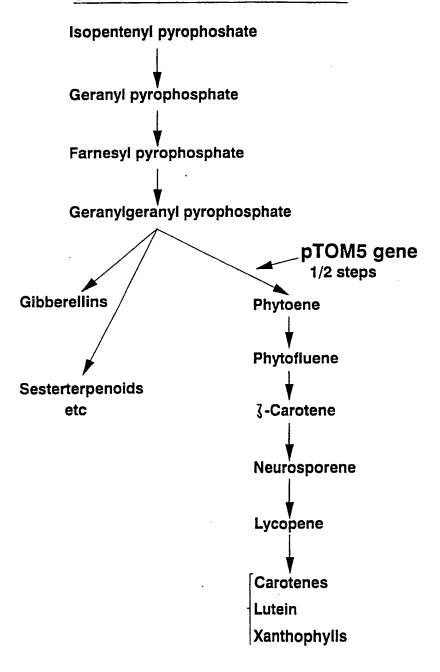
Oligonucleotides used in PCR reactions with pTOM5 to synthesise BamHI-KpnI fragments for cloning into pJR1

Oligo	5'	3'
T5AS-1	**** GGGGG <u>GGATCC</u> TTTGCCTGTCTGTGGTCTTTTTATAATCTT BamHI	
T5AS-2	*** GCAAG <u>GGTACC</u> TTGGTTGTCTAAAACAATTCCAAGATTTTTGTC Kpn I	
T5AS-3	*** CGGACAA <u>GGTACC</u> ATCGAGCATGTCAAATGGCCGCCCATTG Kpn I	
T5AS-4	** * TTTTTTT <u>GGTACC</u> GAAGTCTCTCAAGATAATAAGTGAATACAC Kpn [

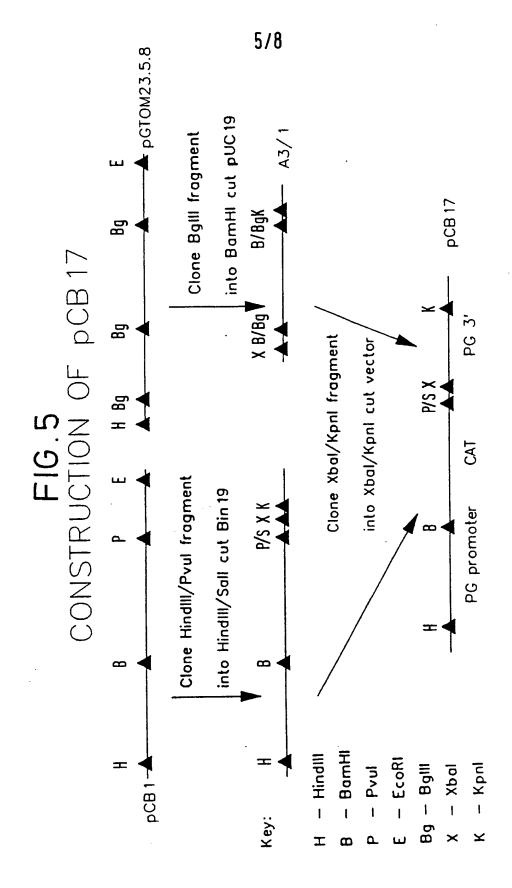
* - base change from pTOM5 sequence

FIG. 4

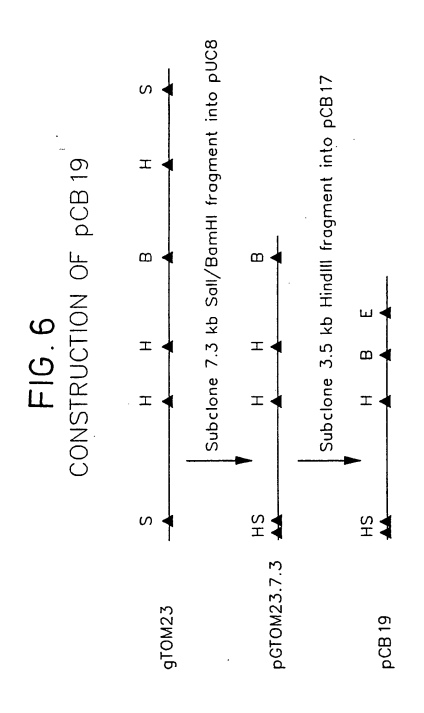
CAROTENOID BIOSYNTHESIS



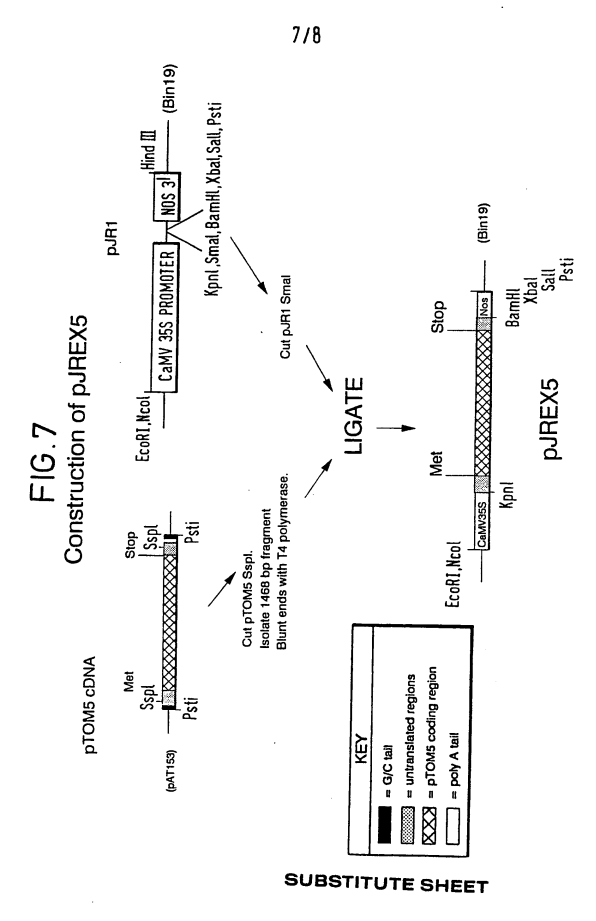
PCT/GB90/01924

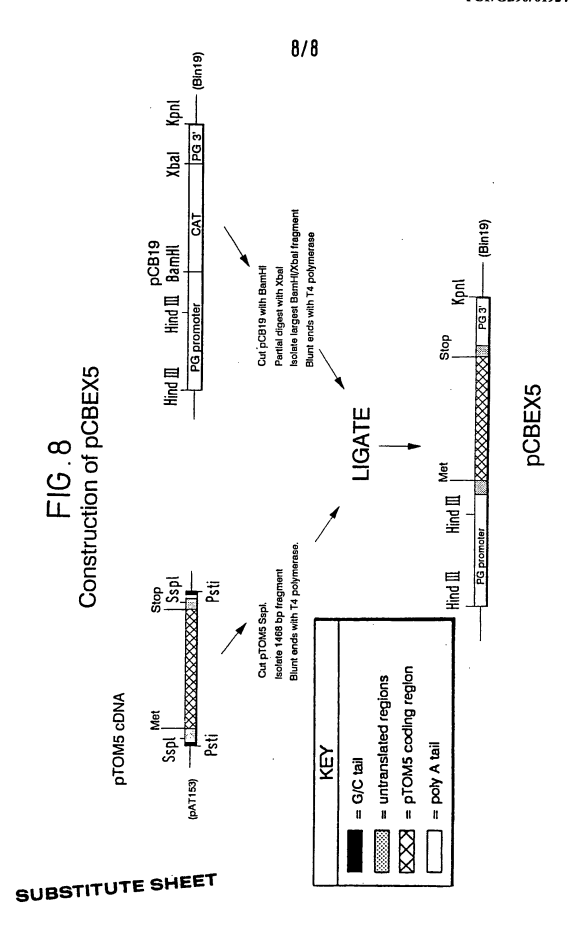


SUBSTITUTE SHEET



Key: S - Sall; H - HindIII; B - BamHI; E - EcoRI





INTERNATIONAL SEARCH REPORT

International Application No PCT/GB 90/01924

		themational Application No 1 C17	35 30/01324	
	IFICATION OF SUBJECT MATTER (it several classif		·	
	to International Patent Classification (IPC) or to both Nati		- 10 - 5/00	
IPC ⁵ :	C 12 N 15/52, 15/29, 15/82 C 12 P 23/00	, A UI H 1/00, 5/00	, C 12 P 5/02	
II. FIELDS	SEARCHED			
	Minimum Documen			
Classificatio	n System	Classification Symbols		
IPC ⁵	C 12 N, A 01 H, C 1	2 P		
	Documentation Searched other t to the Extent that such Documents	han Minimum Documentation are Included in the Fields Searched *		
III. DOCU	MENTS CONSIDERED TO BE RELEVANT			
Category *	Citation of Document, 11 with indication, where app		Relevant to Claim No. 13	
х	WO, A, 88/09334 (CALGENE see page 6, line 32) 1 December 1988,	7,9,12, 14-16,20, 21	
Y	Plant Molecular Biology, 1989, Kluwer Academic W. Schuch et al.: "Commanipulation of gene tomato fruit ripening see page 306, left-hallast 9 lines	c Publ. (BE) ontrol and expression during g", pages 303-311,	1-6,8,11, 13-15,20, 21	
Y	Nucleic Acids Research, v 1987, IRL Press Ltd (J. Ray et al.: "Seque a ripening related cl page 10587, see the v	(Oxford, GB) ence of pTOM5, DNA from tomato",	1-6,8,11, 13-15,20, 21	
Y	Plant Molecular Biology, 1989, Kluwer Academic J.N.M. Mol et al.: "C tion of floral pigmer pages 287-294, see pa	Publ. (BE) Senetic manipula- Station genes",	7,9,10	
"A" doct cont cont cont cont cont cont cont c	categories of cited documents: 19 Iment defining the general state of the art which is not lidered to be of particular relevance or document but published on or after the international of active Iment which may throw doubts on priority claim(s) or this cited to establish the publication date of another ion or other special reason (as specified) Iment referring to an oral disclosure, use, exhibition or Iment multiplished prior to the international filing date but than the priority date claimed	"T" later document published after to or priority date and not in conflicited to understand the principle invention. "X" document of particular relevant cannot be considered novel or involve an inventive step. "Y" document of particular relevant cannot be considered to involve document is combined with one ments, such combination being in the art. "&" document member of the same g	ct with the application but or theory underlying the ce; the claimed invention cannot be considered to ce; the claimed invention an inventive step when the or more other such docubelous to a person skilled patent family	
Date of the	Actual Completion of the International Search	Date of Mailing of this International Se	arch Report	
6th	March 1991	19.04.91	1 11	
	al Searching Authority	Signature of Authorized Officer	in NETO	
EUROPEAN PATENT OFFICE				

111. 00	CUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET	")
Calegory *	Citation of Document, 11 with indication, where appropriate, of the relevant passages	Relevant to Claim No.
Y	The Plant Cell, vol. 1, no. 2, 1989, American Soc. of Plant Physiologists J.O. Narita et al.: "Tomato hydro- xymethylglutaryl-CoA reductase is required early in fruit development but not during ripening", pages 181-190, see page 185, right-hand column, last paragraph; page 186 - page 187, left-hand column	7,9,10
Y	EP, A, 0271988 (IMPERIAL CHEMICAL IND.) 22 June 1988, see the whole document	5,6,10,22
A	EP, A, 0341885 (IMPERIAL CHEMICAL IND.) 15 November 1989, see page 2, column 2, lines 12-17, lines 39-45	5,6,10,16, 22
A	Plant Molecular Biology, vol. 12, 1989, Kluwer Academic Publ. (BE) J. Knapp et al.: "Organization and expression of polygalacturonase and other ripening related genes in Ailsa Craig "neverripe" and "ripening inhibitor" tomato mutants" pages 105-116, see page 106, left- hand column, pages 108-109, page 112, right-hand column; page 114, right-hand column, lines 1-3	7,9,10
A	Theor.Appl. Genet, vol. 76, no. 2, 1988, TAG Springer Verlag M. Mutschler et al.: "Changes in ripening-related processes in tomato conditioned by the alc mutant", pages 285-292, see page 287, right- hand column, page 289, right-hand column; table 4	7,9,10
A	Mol. Gen. Genet, vol. 216, april 1989, MGG Springer Verlag G.A. Armstrong et al.: "Nucleotide sequence, organization, and nature of the protein products of the caro- tenoid biosyntheses gene cluster of Rhodobacter capsulatus", pages 254-268, see figures 1,3	7,9
E	EP, A, 0393690 (KIRIN BEER) 24 October 1990, see page 3, lines 34- 46; page 9, lines 41,42, line 55 - page 10, line 8; claim 7	7,9,12
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Form PCT/ISA 210(extra sheet) (January 1985)

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

GB 9001924 SA 42594

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 12/04/91

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FORM P0479

For more details about this annex : see Official Journal of the European Patent Office, No. 12/82